# SUBSTITUTED 6-BENZYL-4-OXOPYRIMIDINES, PROCESS FOR THEIR PREPARATION AND PHARMACEUTICAL COMPOSITIONS CONTAINING THEM

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The present invention is concerned with compounds which inhibit the reverse transcriptase encoded by human immunodeficiency virus (HIV) or pharmaceutically acceptable salts thereof and are of value in the prevention of infection by HIV, the treatment of infection by HIV and the treatment of the resulting acquired immune deficiency syndrome (AIDS). It also relates to pharmaceutical compositions containing the compounds and to a method of use of the present compounds and other agents for the treatment of AIDS arid viral infection by HIV.

#### BACKGROUND OF THE INVENTION

A retrovirus designated human immunodeficiency virus (HIV) is the etiological agent of the complex disease that includes progressive destruction of the immune system (acquired immune deficiency syndrome; AIDS) and degeneration of the central and peripheral nervous system.

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Currently available drugs for AIDS therapy are divided into two groups: those that prevent infection of target cells [nucleoside (NRTIs) and non-nucleoside reverse transcriptase inhibitors (NNRTIs)], and those that prevent HIV-1-infected cells from yielding infectious viruses (protease inhibitors). Monotherapy with antiretroviral agents has shown limited effects, very likely due to the interplay of phenomena such as: high viral loads and multiplication rates of HIV, incomplete inhibition of viral replication and emergence of drug resistant mutants. For this reason, combination therapies with two or more drugs have been proposed for a more effective treatment of AIDS. Potent suppression of HIV replication over prolonged periods has been accomplished with regimens including reverse transcriptase and protease inhibitors, although on stopping therapies viraemia has rapidly reappeared. In the attempt to obtain better results, research is now focused on exploiting new targets and enhancing the activity of "old" drugs. Among the latter, NNRTs possibly endowed with better pharmacokinetic profiles, capability to inhibit clinically relevant mutants and, hopefully, to minimize HIV multiplication are being pursued.

Compounds of the present invention are dihydro-alkyloxy-benzyl-oxopyrimidines (DABOs) which potently inhibit HIV multiplication targeting reverse transcriptase without bioactivation.

#### BRIEF DESCRIPTION OF THE INVENTION

Novel compounds of formula A:

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as herein defined, are disclosed. These compounds are useful in the inhibition of HIV reverse transcriptase, the prevention of infection by HIV, the treatment of infection by HIV and in the treatment of AIDS, either as compounds, pharmaceutically acceptable salts (when appropriate), pharmaceutical composition ingredients, whether or not in combination with other antivirals, anti-infectives, immunomodulators, antibiotics or vaccines. Methods of treating AIDS, methods of preventing infection by HIV, and methods of treating infection by HIV are also disclosed.

## DETAILED DESCRIPTION OF THE INVENTION AND PREFERRED EMBODIMENTS

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This invention is concerned with the compounds of formula A described below, combinations thereof, or pharmaceutically acceptable salts thereof, in the inhibition of HIV reverse transcriptase, the prevention or treatment of infection by HIV and in the treatment of

the resulting acquired immune deficiency syndrome (AIDS). The compounds of this invention include those with structural formula A:

$$R_4$$
 $R_5$ 
 $R_4$ 
 $R_2$ 
 $R_3$ 
 $R_4$ 
 $R_3$ 
 $R_4$ 
 $R_3$ 
 $R_4$ 
 $R_4$ 
 $R_5$ 
 $R_4$ 
 $R_5$ 
 $R_5$ 
 $R_5$ 
 $R_5$ 
 $R_5$ 
 $R_5$ 
 $R_5$ 
 $R_5$ 
 $R_6$ 
 $R_7$ 

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X is -O, -CH<sub>2</sub>, -CHK (wherein K is -H, -C<sub>1.4</sub> alkyl, -C<sub>3.6</sub>Cycloalkyl), -S, -NK (wherein K is -H, -Cl<sub>1.4</sub> alkyl, -C<sub>3.6</sub> cycloalkyl), -aryl, -arylalkyl;

R is
-H, -C<sub>1.4</sub>alkyl (containing one or more of heteroatoms like 0, S, N), -C<sub>3.6</sub>
cycloalkyl (containing one or more of heteroatoms like 0, S, N), -aryl, -arylakl,
heterocycle;

Y is -H, -C<sub>1.4</sub>alkyl, -C<sub>3.6</sub>cycloalkyl;

15 Z is -H, -C<sub>1-1</sub>alkyl, -C<sub>3-6</sub>cycloalkyl;

R<sub>1</sub> is -H, -C<sub>1-1</sub>alkyl, -halogen, -NO<sub>2</sub>, -OW (wherein W is -H, -CH<sub>3</sub>, aryl), -SW (wherein W is -H, -CH<sub>3</sub>, -aryl);

20 R<sub>2</sub> is -H, -C<sub>1.4</sub>alkyl, -halogen, -NO<sub>2</sub>, (wherein W is -H, -CH<sub>3</sub>, -aryl); -SW (wherein W is -H, -CH<sub>3</sub>, -aryl);

R<sub>3</sub> is -H, -C<sub>1.4</sub>alkyl, -halogen, -NO<sub>2</sub>, -OW (wherein W is -H, -CH<sub>3</sub>, -aryl); -SW (wherein W is -H, -CH<sub>3</sub>, -aryl)

R<sub>4</sub> is -H, -C<sub>1.4</sub>alkyl, -halogen, -NO<sub>2</sub>, -OW (wherein W is -H, -CH<sub>3</sub>, -aryl); -SW (wherein W is -H, -CH<sub>3</sub>, -aryl)

R<sub>3</sub> is
-H<sub>1</sub>, -C<sub>1,4</sub>alkyl, -halogen, -NO<sub>2</sub>, -OW (wherein W is -H<sub>1</sub>, -CH<sub>3</sub>, -aryl), -SW (wherein W is -H<sub>1</sub>, -CH<sub>3</sub>, -aryl);

- pharmaceutically acceptable salts or soluble derivatives thereof;
- preparation process of derivatives thereof;

- a method of preventing infection of HIV, or of treating infection by HIV or of treating AIDS, comprising administering to a mammal an effective amount of compounds claimed;
- a pharmaceutical composition useful for inhibiting HIV reverse transcriptase, comprising an effective amount of compounds claimed, and a pharmaceutically acceptable carrier;
- a pharmaceutical composition useful for preventing or treating infection of HIV or for treating AIDS, comprising an effective amount of compounds claimed, and a pharmaceutically acceptable carrier.

The most preferred compounds of this invention are those of table 1.

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The compounds of the present invention may have asymmetric centers and occur as racemates, racemic mixtures, individual diastereomers, or enantiomers, with all isomeric forms being included in the present invention.

When any variable occurs more than one time in any constituent or in formula A of this invention, its definition on each occurrence is independent of its definition at every other occurrence. Also, combinations of substituents and/or variables are permissible only if such combinations result in stable compounds.

As used herein except where noted, "alkyl" is intended to include both branched- and straight-chain saturated aliphatic hydrocarbon groups having the specified number of carbon atoms; "Halogen" or "Hal" as used herein, means fluoro, chloro, bromo and iodo.

As used herein, with exceptions as noted, "aryl" is intended to mean any stable monocyclic, bicyclic or tricyclic carbon ring of up to 7 members in each ring, wherein at least one ring is aromatic. Examples of such aryl elements include phenyl, naphthyl, tetrahydronaphthyl, biphenyl.

The term heterocycle or heterocyclic, as used herein except where noted represents a stable 5- to 7-membered monocyclic or stable 8- to 11 -membered bicyclic heterocyclic ring which is either saturated or unsaturated, and which consists of carbon atoms and from one to three heteroatoms selected from the group consisting of N, 0 and S; and wherein the nitrogen and sulfur heteroatoms may optionally be oxidized, and the nitrogen heteroatom may optionally be quaternized, and including any bicyclic group in which any of the above-defined heterocyclic rings is fused to a benzene ring. The heterocyclic ring may be attached at any heteroatom or carbon atom which results in the creation of a stable structure.

The pharmaceutically-acceptable salts of the novel compounds of this invention that are capable of salt formation (in the form of water- or oil- soluble or dispersible products) include the conventional non-toxic salts or the quaternary ammonium salts of these compounds, which are formed, e.g.; from inorganic or organic acids or bases.

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In preferred embodiments, a compound of the present invention is administered in combination or alternation with AZT, D4T, FTC (2'.3'-dideoxy-3'-thia-5-fluorocytidine); 3TC (Epivir, Glaxo Wellcome, Inc.), AZDU (3'-Azido-2',3'-dideoxyuridine); 141W94 (amprenavir. GlaxoWellcome, Inc.); Viramune (nevirapine). Rescriptor (delavirdine); or DMP-266 (efavirenz). Other examples of antiviral agents that can be used in combination or alternation with the compounds disclosed herein for HIV therapy include DDI, DDC, Delaviridine, β-LddA, β-L-3'-azido-d5FC, carbovir, acyclovir, interferon, stavudine, CS-92 (3'-azido-2',3'-dideoxy-5-methyl-cytidine), 3'-azido nucleosides, and β-D-dioxolane nucleosides such as β-D-dioxolanylguanine (DXG), β-D-dioxolanyl-2,6-diaminopurine (DAPD), and β-D-dioxolanyl-6-chloropurine (ACP).

Preferred protease inhibitors include indinavir ({1(1,S,2R),5(S)]-2,3,5-trideoxy-N-(2.3-dihydro-2-hydroxy-1H-inden-1-yl)-5-[2-[[(1,1-dimethylethyl)amino]carbonyl]-4-(3-pyridinylmethyl)-1-piperazinyl]-2-(phenylmethyl)-D-erythro-pentoamide sulfate; Merck), nelfinavir (Agouron), ritonavir (Abbot), and saquinavir (Invirase; Roche).

Nonlimiting examples of other compounds that can be administered in combination or alternation with the compounds of the present invention to augment the properties of the drug on administration include abacavir: (1S,4R)-4-[2-amino-6-cyclopropyl-amino)-9H-purin-9-yl]-2-cyclopentene-1-methanol succinate (1592U89, a carbovir analog; Glaxo Wellcome); zidovudine: AZT, 3'-azido-3'-deoxythymidine (Glaxo Wellcome); BILA 1906: N-{1S-[[[3-[2S-{(1,1-dimethylethyl)amino]carbonyl}-4R-]3-pyridinylmethyl)thio]-1-piperidinyl]-2R-hydroxy-1S-(phenylmethyl)propyl]amino]carbonyl]-2-methylpropyl}-2-quinolinecarboxamide (Bio Mega/Boehringer-Ingelheim); BILA 2185: N-(1,1-dimethylethyl)-1-[2S-[[2-2,6-dimethylphenoxy)-1-oxoethyl]amino]-2R-hydroxy-4-phenylbutyl]4R-pyridinylthio)-2-piperidinecarboxamide (Bio Mega/Boehringer-Ingelheim); BM+51.0836:triazoloisoindolinone derivative; BMS 186,318: aminodiol derivative HIV-1 protease inhibitor (Bristol-Myers-Squibb); d4API: 9-[2,5-dihydro-5-(phosphonomethoxy)-2-furanel]adenine (Gilead); stavudine: d4T, 2',3'-didehydro-3'-deoxythymidine (Bristol-Myers-Squibb); efavirenz: DMP-266, a 1,4-dihydro-2H-3, 1-benzoxazin-2-one; HBY097: S-4-

isopropoxycarbonyl-6-methoxy-3-(methylthio-methyl)-3,4-dihydroquinoxalin-2(1H)-thione; HEPT: 1-[(2-hydroxyethoxy)methyl]6-(phenylthio)thymine; KNI-272: (2S,3S)-3-amino-2hydroxy-4-phenylbutyric acid-containing tripeptide; L-697,593; 5-ethyl-6-methyl-3-(2phthalimido-ethyl)pyridin-2(1H)-one; L-735,524: hydroxy-aminopentane amide HIV-1 protease inhibitor (Merck); L-697,661: 3-{[(-4,7-dichloro-1,3-benzoxazol-2yl)methyl]amino}-5-ethyl-6-methylpyridin-2(1H)-one; L-FDDC: (-)-\beta-L-5-fluoro-2',3'dideoxycytidine; L-FDOC: (-)-β-L-5-fluoro-dioxolane cytosine; 6-benzyl-1-ethoxymethyl-5isopropyluracil (I-EBU; Triangle/Mitsubishi); nevirapine: 11-cyclopropyl-5,11-dihydro-4methyl-6H-dipyridol[3,2-b:2',3'-e]diazepin-6-one (Boehringer-Ingelheim); PFA: phosphonoformate (foscarnet; Astra); PMEA: 9-(2-phosphonylmethoxyethyl) adenine 10 (Gilead); PMPA: (R)-9-(2-phosphonyl-methoxypropyl)adenine (Gilead); Ro 31-8959: hydroxythethylamine derivative HIV-1 protease inhibitor (Roche); RPI-3121: peptidyl protease inhibitor, 1-[(3s)-3-(n-alpha-benzyloxycarbonyl)-1-asparginyl)-amino-2-hydroxy-4phenylbutyryl]-n-tert-butyl-1-proline amide; 2720: 6-chloro-3,3-dimethyl-4-(isopropenyloxycarbonyl)-3,4-dihydro-quinoxalin-2(1H)thione; SC-52151: hydroxyethylurea isostere protease inhibitor (Searle); SC-55389A: hydroxyethyl-urea isostere protease inhibitor (Searle); TIBO R82150: (+)-(5S)-4,5,6,7-tetrahydro-5-methyl-6-(3-methyl-2butenyl)imidazo[4,5,1-jk][1,4]-benzodiazepin-2(1H)-thione (Janssen); TIBO 82913: (+)-(5S)-[1,4]benzodiazepin-2(1H)-thione (Janssen); TSAO-m3T:[2',5'-bis-O-(tert-20 butyldimethylsilyl)-3'-spiro-5'-(4'-amino-1',2'-oxathiole-2',2'-dioxide)]-  $\beta$ -Dpentofuranosyl-N3-methylthymine; U90152: 1-[3-[(1-methylethyl)-amino]2-pyridinyl]-4-[[5-[(methylsulphonyl)-amino]-1H-indol-2yl]carbonyl]piperazine; UC: thiocarboxanilide derivatives (Uniroyal); UC-781 =N-[4-chloro-3-(3-methyl-2-butenyloxy)phenyl]-2-methyl-3furancarbothioamide; UC-82 = N-[4-chloro-3-(3-methyl-2-butenyloxy)phenyl]-2-methyl-3-25 thiophenecarbothioamide; VB 11,328: hydroxyethylsulphonamide protease inhibitor (Vertex); VX-478: amprenavir, 141W94, hydroxyethylsulphonamide protease inhibitor (Vertex/Glaxo Wellcome); XM 323: cyclic urea protease inhibitor (Dupont Merck), delaviridine (Pharmacia Upjohn), famciclovir, gancyclovir, and penciclovir. In another embodiment,a compound of the present invention is administered in combination with 30

LG1350, which has the following structure.

# Preparation Of Methyl Arylacetylalkylacetates

#### SCHEME A

OH

$$R_1$$
 $R_2$ 
 $R_3$ 
 $R_4$ 
 $R_2$ 
 $R_3$ 
 $R_4$ 
 $R_4$ 
 $R_5$ 
 $R_5$ 

Anhydrous pyridine (400 mmoles, 32.5 ml) was added with stirring under nitrogen atmosphere into an ice-cooled solution of 2,2-dimethyl-1,3-dioxane-4,6-dione (Meldrurm's acid) (165 mmoles, 23.75 g) in anhydrous dichloromethane (50 ml). The resulting solution was treated, over a 2 h period at 0°C under nitrogen atmosphere, with a solution of crude arylacetyl chloride in anhydrous dichloromethane (50 ml). Arylacetyl chloride was prepared before use by refluxing the proper arylacetic acid (43.2 mmoles) with thionyl chloride (21.3 ml) under nitrogen atmosphere for 2 h. Then, the mixture was stirred for 2 h at room temperature, poured into crushed ice and treated with 2N HCl (100 ml). The organic layer was separated and the aqueous solution was extracted twice with dichloromethane (25 ml). The organic phase and the extracts were combined, washed with brine, dried and evaporated. The solid residue was dissolved in anhydrous methanol (250 ml) and the solution was refluxed for 20 h. After cooling, metal sodium (0.16 g-atoms, 3.68 g) was carefully added and the mixture was stirred until dissolution was complete. Alkyl halide (160 mmoles) was dropped into the solution and the resulting mixture was heated at reflux for 4-12 h. After cooling, the solvent was removed and the residue treated with water (200 ml) and extracted with chloroform (3 x 100 ml). The organic layer was washed with brine (2 x 100 ml), dried and evaporated to give the desired compound, which was purified by passing through a silica gel column (chloroform as eluent).

In the above reaction, arylacetic acid (Scheme "A") or arylacetyl chloride can be replaced with the corresponding 1-arylacetylimidazolide (Scheme "B") or with arylacetylethoxycarbonylanhydride, whereas the Meldrum's acid can be replaced with ethyl acetylacetate, ethyl alkylmalonate or ethyl alkylmalonate potassium salt, to give the proper ethyl arylacetylalkylacetates in high yields.

### Preparation Of Compounds (I) With X = O (Scheme A).

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The proper methyl arylacetylalkylacetate (10 mmoles) in methanol (50 ml) was added to a well-stirred suspension of O-methylisourea hydrogen sulphate (15 mmoles, 2.58 g) and calcium hydroxide (16 mmoles, 1.18 g) in water (50 ml). The resulting mixture was stirred at room temperature for 72 h, then concentrated, made acid (pH 5) with 0.5N acetic acid and extracted with ethyl acetate (3 x 50 ml). The combined organic extracts were washed with brine (100 ml), dried and evaporated to dryness. The residue was purified by crystallization

from the proper solvent yielding pure 5-alkyl-6-benzyl-3,4-dihydro-2-methoxypyrimidin-4-one. This compound was then refluxed with the proper potassium alkoxide (100 mmoles of potassium metal in 20-30 ml of alcohol freshly distilled on sodium metal) under nitrogen atmosphere until starting material disappeared at the TLC control. After cooling, the mixture was concentrated, made acid (pH 5) with 0.5N acetic acid and extracted with ethyl acetate (3 x 50 ml). The combined extracts were washed once with brine (100 ml), dried and evaporated to give the required 2-alkoxy-5-alkyl-6-benzyl-3,4-dihydropyrimidin-4-one derivative, which was recrystallized from a suitable solvent or purified by column chromatography (silica gel; ethyl acetate:chloroform 1:1). Physical and chemical data of representative compounds of the invention are reported in table 1; cytotoxicity and anti-HIV-1 activity data are reported in table 2.

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## Preparation Of Compounds (I) With X = S

#### SCHEME B

The proper ethyl arylacetylalkylacetate (31.5 mmoles) was successively added to a stirred solution of sodium metal (0.063 g-atoms) in 50 mL of absolute ethanol (50 ml) thiourea (43 mmoles). The mixture was heated while stirring at reflux for 5 h. After cooling, the solvent was distilled *in vacuo* at 40-50°C until dryness and the residue was dissolved in water (200 mL) and made acid (pH 5) with 0.5N acetic acid. The resulting precipitate (the crude 2-thiouracil derivative) was filtered under reduced pressure, washed with diethyl ether, vacuum dried at 80°C for 12 h and then crystallized from the proper solvent.

Then, according to method A, iodomethane (8 mmoles, 1.13 g) was added to a suspension containing the proper 2-thiouracil derivative (4 mmoles) in anhydrous N,N-dimethylformamide (2 ml), and the resulting mixture was stirred at room temperature until the starting material disappeared at the TLC control (silica gel; n-hexane: ethyl acetate: methanol 12:3:1). Then the reaction content was poured on cold water (100 mL) and extracted with ethyl acetate (3 x 50 ml). The organic layers were collected, washed with a sodium thiosulfate solution (100 ml), brine (3 x 50 ml), dried and evaporated to furnish the crude 5-alkyl-6-benzyl-3,4-dihydro-2-methylthiopyrimidin-4-one (2) as a solid purified by crystallization.

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Alternatively, according to methods B and C, potassium carbonate (4.2 mmoles) and the proper alkyl halide (4.4 mmoles) were added to a suspension containing 2-thiouracil derivative (4 mmoles) in anhydrous N,N-dimethylformamide (2 ml). The resulting mixture was stirred at room temperature (method B) or at 80°C (method C) until starting material disappeared at the TLC control (silica gel; n-hexane:ethyl acetate:methanol 12:3:1). Then the reaction content was poured on cold water (200 mL), made acid (pH 5) with 0.5N acetic acid and extracted with ethyl acetate (3 x 50 ml). The organic layers were collected, washed with a sodium thiosulfate solution (100 ml), brine (100 ml), dried and evaporated to furnish 5-alkyl-6-benzyl-3,4-dihydro-2-methylthiopyrimidin-4-ones (3) and (4) as crude material which was then purified by column chromatography on silica gel (eluent: n-hexane:ethyl acetate:methanol 12:3:1) followed by crystallization. Physical and chemical data of representative compounds of the invention are reported in table 1. Cytotoxicity and anti-HIV-1 activity in vitro are reported in table 2.

## Preparation Of Compounds (I) With X = NK

#### SCHEME C

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Title derivatives were prepared according to the procedure described for the synthesis of compounds with X = S(I), using ethyl arylacetylalkylacetates and guanidine [2-amino-6benzylpyrimidin-4-ones (5)] as starting materials. 2-Alkylaminoderivatives (6) were synthesized by heating the previously reported 5-alkyl-6-benzyl-3,4-dihydro-2-methylthio pyrimidin-4-ones with 20-30 ml of proper amine in a sealed tube at 170°C for 24 h. Physical and chemical data of some compounds (6) are reported in table 1. Cytotoxicity and anti-HIV-I activity in vitro are reported in table 2. The compounds of the present invention are useful in the inhibition of HIV reverse transcriptase, the prevention or treatment of infection by the human immunodeficiency virus (HIV) and the treatment of consequent pathological conditions such as AIDS. Treating AIDS or preventing or treating infection by HIV is defined as including, but not limited to, treating a wide range of states of HIV infection: AIDS, ARC (AIDS related complex), both symptomatic and asymptomatic, and actual or potential exposure to HIV. For example, the compounds of this invention are useful in treating infection by HIV after suspected past exposure to HIV by, e.g., blood transfusion, organ transplant, exchange of body fluids, bites, accidental needle stick, or exposure to patient blood during surgery.

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The compounds of this invention are also useful in the preparation and execution of screening for antiviral compounds. For example, the compounds of this invention are useful for isolating enzyme mutants, which are excellent screening tools for more powerful antiviral compounds. Furthermore, the compounds of this invention are useful in establishing or determining the binding site of other antiviral to HIV reverse transcriptase e.g., by competitive inhibition. Thus the compounds of this invention are commercial products to be sold for these purposes. For inhibition of HIV reverse transcriptase, the prevention or treatment of infection by HIV and the treatment of AIDS or ARC, the compounds of the present invention may be administered orally, parenterally (including subcutaneous injections, intravenous, intramuscular, intrasternal injection or infusion techniques), by inhalation spray, or rectally, in dosage unit formulations containing conventional non toxic pharmaceutically-acceptable carriers, adjuvants and vehicles. Thus, in accordance with the present invention there is further provided a method of treating and a pharmaceutical composition for treating HIV infection and AIDS. The treatment involves administering to a patient in need of such treatment a pharmaceutical composition comprising a pharmaceutical carrier and a therapeutically effective amount of a compound of the present invention. These pharmaceutical compositions may be in the form of orally administrable suspensions or tablets; nasal sprays; sterile injectable preparations, for example, as sterile injectable aqueous or oleagenous suspensions or suppositories.

When administered orally as a suspension, these compositions are prepared according to techniques well-known in the art of pharmaceutical formulation and may contain microcrystalline cellulose for imparting bulk, alginic acid or sodium alginate as a suspending agent, methylcellulose as a viscosity enhancer, and sweetners/flavoring agents known in the art. As immediate release tablets, these compositions may contain microcrystalline cellulose, dicalcium phosphate, starch, magnesium stearate and lactose and/or other excipients, binders, extenders, disintegrants, diluents and lubricants known in the art.

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When administered by nasal aerosol or inhalation, these compositions are prepared according to techniques well-known in the art of pharmaceutical formulation and may be prepared as solutions in saline, employing benzyl alcohol or other suitable preservatives, absorption promoters to enhance bioavailability, fluorocarbons, and/or other solubilizing or dispersing agents known in the art.

The injectable solutions or suspensions may be formulated according to known art, using suitable non toxic, parenterally acceptable diluents or solvents, such as mannitol, 1,3-butanediol, water, Ringer's solution or isotonic sodium chloride solution, or suitable dispersing or wetting and suspending agents, such as sterile, bland, fixed oils, including synthetic mono- or diglycerides, and fatty acids, including oleic acid.

When rectally administered in the form of suppositories, these compositions may be prepared by mixing the drug with a suitable non-irritating excipient; such as cocoa buffer, synthetic glyceride, esters or polyethylene glycols, which are solid at ordinary temperatures, but liquidity and/or dissolve in the rectal cavity to release the drug.

The compounds of this invention can be administered orally to humans in a dosage range of 1 to 75 mg/kg body weight. One preferred dosage range is 1 to 50 mg/kg body weight orally. Another preferred dosage range is 5 to 75 mg/kg body weight orally. It will be understood, however, that the specific dose level and frequency of dosage for any particular patient may be varied and will depend upon a variety of factors including the activity of the specific compound employed, the metabolic stability and length of action of that compound, the age, body weight, general health, sex, diet, mode and time of administration, rate of

excretion, drug combination, the severity of the particular condition, and the host undergoing therapy.

The present invention is also directed to combinations of the HIV reverse transcriptase inhibitor compounds with one or more agents useful in the treatment of AIDS. The compounds of this invention can be administered in combination with other compounds that are HIV reverse transcriptase inhibitors, and/or with compounds that are HIV protease inhibitors. When used in a combination treatment with compounds of the instant invention, dosage levels of HIV protease inhibitors of the order of 1 to 25 or 50 grams-per-day are useful in the treatment or prevention of the above-indicated conditions, with oral doses two-to-five time higher. For example, infection by HIV is effectively treated by the administration of from 5 to 25 milligrams of the HIV protease inhibitor per kilogram of body weight from one to three times per day.

It will be understood, however, that the specific dose level and frequency of dosage for any particular patient may be varied and will depend upon a variety of factors including the activity of the specific compound employed, the metabolic stability and length of action of that compound, the age, body weight, general health, sex, diet, mode and time of administration, rate of excretion, drug combination, the severity of the particular condition, and the host undergoing therapy. Dosages of HIV reverse transcriptase inhibitors, when used in a combination treatment with compounds of the present invention, are comparable to those dosages specified above for the present compounds. It will be understood that the scope of combinations of the compounds of this invention with AIDS antivirals includes any combination with any pharmaceutical composition useful for the treatment of AIDS.

#### ANTIVIRAL ASSAY PROCEDURES

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Compounds. Compounds were solubilized in DMSO at 200 mM and then diluted into culture medium.

Cells and viruses. MT-4, C8166, H9/IIIB and CEM cells were grown at 37 °C in a 5% CO<sub>2</sub> atmosphere in RPMI 1640 medium, supplemented with 10% fetal calf serum (FCS), 100 IU/mL penicillin and 100 µg/mL streptomycin. Cell cultures were checked periodically for the absence of mycoplasma contamination with a MycoTect Kit (Gibco). Human

immunodeficiency virus type-1 (HIV-1, III<sub>B</sub> strain) was obtained from supernatants of persistently infected H9/III<sub>B</sub> cells. HIV-1 stock solution had a titres of  $4.5 \times 10^6 50\%$  cell culture infectious dose (CCID<sub>50</sub>)/ml.

HIV titration. Titration of HIV was performed in C8166 cells by the standard limiting dilution method (dilution 1:2, four replica wells per dilution) in 96-well plates. The infectious virus titre was determined by light microscope scoring of cytopathicity after 4 days of incubation and the virus titres were expressed as CCID<sub>50</sub>/mL.

Anti-HIV assays. Activity of the compounds against HIV-1 and HIV-2 multiplication in acutely infected cells was based on the inhibition of virus-induced cytopathicity in MT-4 and C8166 cells, respectively. Briefly, 50 µL of culture medium containing lxl0<sup>4</sup> cells were added to each well of flat-bottom microtiter trays containing 50 µl of culture medium with or without various concentrations of the test compounds. Then 20 µL of an HIV suspension containing 100 CCID<sub>50</sub> were added. After a 4-day incubation at 37 °C, the number of viable cells was determined by the 3-(4,5-dimethylthiazol-1-yl)-2,5-diphenyltetrazolium bromide (MTT) method. Cytotoxicity of the compounds was evaluated in parallel with their antiviral activity. It was based on the viability of mock-infected cells, as monitored by the MTT method.

RT assays. Assays were performed as follows. Briefly, purified rRT was assayed for its RNA-dependent polymerase-associated activity in a 50  $\mu$ L volume containing: 50 mM TrisHCl (pH 7.8), 80 mM KCll, 6mM MgCl2, 1 mM DTT, 0.1 mg/ mL BSA, 0.3 OD<sub>260</sub> unit/mL template:primer [poly(rC)-oligo(dG)12-18] and 10  $\mu$ M [ $^3$ H]dGTP (1 Ci/mmol). After incubation for 30 min at 37 °C, the samples were spotted on glass fiber filters (Whatman GF/A), and the acid-insoluble radioactivity was determined.

#### 25 EXAMPLES

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2-Cyclopentylthio-6-(2,6-difluorophenylmethyl)-3,4-dihydrogyrimidin-4-(3H)-one (MC867). A mixture of 6-(2,6-difluorophenylmethyl)-1,2,3,4-tetrahydro-2-thiopyrimidin-4(3H)-one (0.16 g, 0.65 mmol; prepared as reported in scheme B), cyclopentyl bromide (0.11 g, 0.08 mL., 0.71 mmol) and potassium carbonate (0.09 g, 0.65 mmol) in 1 mL of anhydrous DMF was stirred at room temperature for 24 h. After treatment with cold water (200 mL), the solution was extracted with ethyl acetate (3 x 50 mL). The organic layers were collected, washed with brine (3 x 50 mL), dried and evaporated to furnish crude MC867, which was

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purified by chromatography on silica gel column (eluent: n-hexane/ethyl acetate/methanol 12/3/1).

Yield (%): 45; mp (°C): 168-169; recrystallization solvent: cyclohexane; formula (molecula-weight):  $C_{16}H_{16}F_2N_2OS$  (322.37).

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2-Cyclopenlylthio-6-(2,6-difluorophenylmethyl)-3,4-dihydro-5-methylpyrimidin-4-(3H)-one (MC922).

The synthesis of MC922 was accomplished according to the above reported procedure starting from 6-(2,6-difluorophenylmethyl)-5-methyl-1,2,3,4-tetrahydro-2-thiopyrimidin-4-(3H)-one (see scheme B).

Yield (%): 54; mp (°C): 192-193; recrystallization solvent: cyclohexane; formula (molecular weight): C<sub>17</sub>H<sub>18</sub>F<sub>2</sub>N<sub>2</sub>OS (336.40).

# 2-Cyclopentylthio-6-[l-(2,6-difluorophenyl)ethyl]-3,4-dihydropyrimidin-4-(3H)-one (MC1008)

The synthesis of MC1008 was accomplished according to the above reported procedure starting from 6-[1-(2,6-difluorophenyl)ethyl]-1,2,3,4-tetrahydro-2-thiopyrimidin-4(3H)-one (see scheme B).

Yield (%): 54; mp (°C): 165.5-166.5; recrystallization solvent: cyclohexane; formula (molecular weight):  $C_{17}H_{18}F_2N_2OS$  (336.40).

# 2-Cyclopentylthio-6-[l-(2,6-difluorophenyl)ethyl]-3,4-dihydro-5-methylpyrimidin4(3H)-one (MC1047)

The synthesis of MC1047 was accomplished according to the above reported procedure, starting from 6-[l-(2,6-difluorophenyl)ethyl]-5-methyl-1,2,3,4-tetrahydro-2-thiopyrimidin-4(3H)-one (see scheme B).

Yield (%): 60; mp (°C): 196-197; recrystallization solvent: cyclohexane; formula (molecular weight):  $C_{18}H_{20}F_2N_2OS$  (350.43).

### 6-(2,6-Difluorophenylmethyl)-3,4-dihydro-2-(methylthiomethyl)thiopyrimidin-4-(3H)-one (MC1161)

The synthesis of MC1161 was accomplished according to the above reported procedures, starting from 6-(2,6-difluorophenylmethyl)-1,2,3,4-tetrahydro-2-thiopyrimidin-4(3H)-one (see scheme B) and chloromethyl methyl sulfide.

Yield (%): 72; mp (°C): 159-160; recrystallization solvent: benzene/cyclohexane; formula (molecular weight):  $C_{13}H_{12}F_2N_2OS_2$  (314.37).

6-(2.6-Difluorophenylmethyl)-3,4-dihydro-5-methyl-2-(methylthiomethyl)thiopyrimidin-4(3H)-one (MC1162). 10

The synthesis of MC1162 was accomplished according to the above reported procedure, starting from 6-(2,6-difluorophenylmethyl)-5-methyl-1,2,3,4-tetrahydro-2-thiopyrimidin 4(3H)-one (see scheme B) and chloromethyl methyl sulfide.

Yield (%): 70; mp (°C): 183-184; recrystallization solvent: benzene/cyclohexane; formula (molecular weight):  $C_{14}H_{14}F_2N_2OS_2$  (328.39).

6-(2.6-Difluorophenylmethyl)-3.4-dihydro-5-(1-methylethyl)-2-(methylthiomethyl) thiopyrimidin-4-(3H)-one MC1145).

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The synthesis of MC1145 was accomplished according to the above reported procedure, starting from 6-(2,6-difluorophenylmethyl)-5-(1-methylethyl)-1,2,3,4-tetrahydro-2-20 thiopyrimidin-4(3H)-one (see scheme B) and chloromethyl methyl sulfide. Yield (%): 62; mp (°C): 158.5-160; recrystallization solvent: cyclohexane; formula (molecular weight):  $C_{16}H_{18}F_2N_2OS_2$  (356.45).

2-Cyclopenltylamino-6-(2.6-difluorophenylmethyl)-3,4-dihydropyrimidin-4-(3H)-one 25 (MC1022).

Cyclopentylamine (10 mL) was heated while stirring with 6-(2,6-difluorophenylmethyl)-3,4dihydro-2-methylthiopyrimidin-4-(3H)-one (0.30 g, 1.12 mmol; prepared as reported in scheme B or C) in a sealed tube at 160°C for 10 h. After cooling, the mixture was diluted with water (200 mL) and extracted with ethyl acetate (3 x 50 mL). The organic layers were collected, washed with brine (3 x 50 mL), dried and evaporated to furnish crude MC1022,

which was purified by chromatography on silica get column (eluent: ethyl acetate/chloroform 1/1).

Yield (%): 74; mp (°C): - (oil); formula (molecular weight):  $C_{16}H_{17}F_2N_3O$  (305.33).

- 2-Cyclopentylamino-6-(2.6-difluorophenylmethyl)-3.4-dihydro-5-methylpvrimidin-4-(3H)-5 one (MC1050).
  - The synthesis of MC1050 was accomplished according to the above reported procedure, starting from 6-(2,6-difluorophenylmethyl)-3,4-dihydro-5-methyl-2-methylthiopyrimidirin-4(3H)-one (see scheme B or C).
- Yield (%): 60; mp (°C): 115-117; recrystallization solvent: n-hexane/cyclohexane; formula 10 (molecular weight):  $C_{17}H_{19}F_2N_3O$  (319.35).
  - 2-Cyclopentylamino-6-[1-(2,6-difluorophenyl)ethyl]-3,4-dihydropyrimidin-4-(3H)-one (MC1048).
- The synthesis of MC1048 was accomplished according to the above reported procedure, starting from 6-[1-(2,6-difluorophenyl)ethyl]-3,4-dihydro-2-methylthiopyrimidin-4(3H)-one (see scheme B or C).
  - Yield (%): 48; mp (°C): (oil); formula (molecular weight)  $C_{17}H_{19}F_2N_3O$  (319.35).
- 2-Cyclopentylamino-6-[l-(2.6-difluorophenyl)ethyl]-3,4-dihydro-5-methylpyrimidin-4-(3H)-20 one (MC1129)
  - The synthesis of MC1129 was accomplished according to the above reported procedure, starting from 6-[1-(2,6-difluorophenyl)ethyl]-3,4-dihydro-5-methyl-2-methylthiopyrimidin-4(3H)-one (see scheme B or C).
- Yield (%): 38; mp (°C): (oil); formula (molecular weight):  $C_{18}H_{21}F_2N_3O$  (333.38).
  - 6-(2.6-Difluorophenylmethyl)-3.4-dihydro-2-(4-thiomorpholin-1-yl)pyrimidin-4-(3H)-one (MC1193).
- The synthesis of MC1193 was accomplished according to the above reported procedure, starting from thiomorpholine and 6-(2,6-difluorophenylmethyl)-3,4-dihydro-2-30 methylthiopyrimidin-4(3H)-one (see scheme B or C).

Yield (%): 78; mp (°C): 233-234; recrystallization solvent: acetonitrile; formula (molecular weight): C<sub>15</sub>H<sub>15</sub>F<sub>2</sub>N<sub>3</sub>OS (323.36).

6-(2.6-Difluorophenylmethyl)-3.4-dihydro-2-N.N-dimethylaminopyrimidin-4-(3H)-one (MC1182).

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To a stirred solution of sodium metal (0.14 g, 6.3 mg-atoms) in absolute ethanol (50 mL) 1,1-dimethylguanidine sulfate (1.17 g, 4.3 mmol) and ethyl 4-(2,6-difluorophenyl)acetylacetate (0.76 g, 3.15 mmol) were successively added. The mixture was heated while stirring at reflux for 8 h. After cooling, the solvent was distilled *in vacuo* at 40-50°C until dryness and the residue was dissolved in water (200 mL) and made acid (pH 5) with 0.5N acetic acid. The resulting precipitate (the crude isocytosine derivative) was filtered under reduced pressure, washed with diethyl ether, vacuum dried at 80°C for 12 h and then crystallized from benzene/cyclohexane (see scheme C starting from ethyl 4-(2,6-difluorophenyl)acetylacetate and replacing guanidine hydrochloride with 1,1-dimethylguanidine sulfate).

Yield (%): 88; mp (°C): 210-211; recrystallization solvent: benzene/cyclohexane; formula (molecular weight): C<sub>13</sub>H<sub>13</sub>F<sub>2</sub>N<sub>3</sub>O (265.26).

Table 1. Physical and Chemical Data of MC Compounds

					c ·	ć'	ņ		· .	· ·	so.	s:	s.	×	S	s	s	<u>ર્</u>	ž	S.	v.	v.	SO	SC.	S	S	š	Š	SO.	SO	2	
Formula "	C <sub>IMIN</sub> N <sub>2</sub> O <sub>2</sub>	C <sub>19</sub> H <sub>24</sub> N <sub>2</sub> O <sub>2</sub>	Charano	Carran	CitlinFiN	Chilerin	ChilkN	C,11,10,10	C <sub>15</sub> H <sub>14</sub> N <sub>2</sub> OS	Cultur	CIMITANOS	O'N'E	Culting	C'N'T'U	Cle Han NiOS	C, H, B, N, O	CthHan	Cistingo	C,NII,NO	CISHINNOS	C <sub>15</sub> H <sub>17</sub> N <sub>3</sub> O	CisHiNOS	CISHINCINIOS	C <sub>15</sub> H <sub>1</sub> /CIN <sub>2</sub> OS	C, H, F,	CISHIPFN	Cl. Hww.os	C. HWN, OS	C.N.I.,I.,N.OS	C. II, F. N.OS	CLECK	
% yicld	22	28	2	<u>∞</u>	88	22	æ	20	æ :	65	59	<b>8</b> 2	69	9	<b>L9</b>	<u>5</u>	26	89	<b>Z</b> .	9	88	<b>5</b> ,	74	89	47	74	74	11	68	75	9	;
Recryst. Solvent	Petrol. Ether/dicthyl ether	Petrol. Ether/diethyl ether	Petrol. Ether/diethyl ether	Petrol. Ether/diethyl ether	Petrol. Ether/diethyl ether	Benzene	Cyclohexane/benzene	n-hexane/cyclohexane	n-hexane	n-hexane	n-hexane	Cyclohexane	Cyclohexane	Cyclohexane	n-hexane/cyclohexane	Cyclohexane	n-hexanc	Cyclohexane/benzene	Cyclohexane/benzene	Petrol. Ether/diethyl ether	n-hexane/cyclohexane	Cyclohexane	Cyclohexane	n-hexane/cyclohexane	Cyclohexane	n-hexane	Cyclohexane/henzene	Cyclohexane	Cyclohexane	Cyclohexane	Cyclohexane	
п.р., °С	130-132	132-134	178-181	196-198	87-88	183.5-184.5	181-183	157-158	118-119	96-56	142-143	144-145	168-169	175.5-176.5	118-119	142-144	107.5-108.5	148.0-148.5	127-128	128-130	120-121	66-86	125-126	106-107	76-96	66-86	143-144	128-130	125-126	144-145	122-124	16.1-1.4
5≃	×	=	=	=	Ľ	ب	Ξ	=	Ξ	I	=	Ξ	Ξ	=	×	=	=	=	=	Ξ	I	=	=	=	I	=	=	Ξ	=	: =	: =	=
~	I	I	I	I	Ξ	=	=	Ŧ	=	=	=	=	Ξ	=	=	=	=	=	=	×	I	=	=	=	Ξ	=	=	=	: =	: =	: :	=
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R <sup>2</sup>	=	=	=	I	=	=	=	Ξ	I	=	Ξ	=	=	Ξ	=	=	: =	=	Ź	=	Ξ	ວ	=	=	٠	=	=	: =	: =	: =	: :	=
<u>-</u> ≃	Ξ	=	Ξ	Ξ	ت	Ĺ	=	Ξ	=	=	=	=	=	: =	: W	Ž	Ξ	ć	· =	===	5	=	I	·-	=	: =	Ž	: : :	= č	֓֞֝֟֞֞֝֞֞֓֓֞֓֓֞֞֓֓֓֓֞֓֓֓֓֞֓֓֓֞֞֓֓֡֓֞֓֓֓֞֜֜֡֓֡֡֡֡֡֓֓֡֡֡֡֡֡֡֡	= ;	OMe
~	2.5-Mey-c-hex	4,5-Me,-c-hex	3,5-Me,-c-hex	2.5-Me,-c-hex	Sec-but	c-pent	benzyloxymeth	Sec-but	lso-prop	c-pent	c-hex	lso-prop	c-nent	-Prox	Sechul	C-nent	Sec-but	Sechul	Sec-but	Sec-but	Sec-but	Sec-but	Sec-but	Sec-but	Sec-but	Sec-but	Sec-but	Sea but	Sec-put	Sec-pul	Sec-001	Sec-but
2	Ξ	=	=	Ξ	=	=	=	£	M	Me	ž	ũ	ē	Œ	i =	: =	: =	: =	: =	: =	=	=	=	=	: =	: =	: =	= =	= :	= :	= :	=
>	Ξ	=	=	M	=	=	: =	=	=	=	: =	=	=	: =	: =	: =	: =	= =	: =	: =	: =	: =	: =	: =	: =	: =	: =	= =	= :	= :	= :	=
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Compd.	MC 507	MC 508	MC 512	MC 531	MC 1114	MC 1103	MC 843	MC 796	MC 890	MC 892	MC 808	MC 800	MC 900	MAC 003	MC 20.5	000 JW	MC 900	MC 917	MC 807	MC 863	MC 854	MC 857	MC 850	MC 880	Pas JW	100 OF	MC 003	MC 023	MC 900	MC 868	MC 454	MC 952

Fable 1.		_	Physica	sical and Chemical Data of MC Compounds (continued)	Data of MC C	nodute	nds (co	ntinuc	<del>-</del>				
Compd.	×	>	2	~	R' R <sup>2</sup>		, ×	~	\$2	m.p., °C	Recryst, Solvent	% yield	Formula "
4((.057	·	=	Ξ	Sershut	0	OMe	_	=	=	78-80	n-hexane/ Cyclothexane	11	C.I.II.N.O.S
MC 964	: 🗸	: =	: =	Sec-but	=======================================		OMe	=	=	112-113	Cyclohexane	S	SONETH
MC 1041	· v	: =	: =	Sec-but	=		_	Ξ	=	122-123	Cyclohexane	ž	CialiFiniOS
MC 1042	· 55	=	=	Sec-but	=	Mc	_	=	=	119-120	n-hexane	77	C, Nully No.OS
MC877	: 25	=	=	Me	5	_	_	=	5	237-238	benzene	×	C.I.I.I.C.I.N.O.S
MC878	· v	=	=	iso-prop	=	_	_	=	<u>.</u>	230-231	benzene	≆	CITICINOS
MC886	s	=	=	. Ind-u	= :	_	_	=	ວ	153-154	cyclohexane	62	CISHIP CINIOS
MC885	S	=	=	iso-but	= :	_	_	=	ວ	143.5-144.5	cyclohexane	26	CISH,CLNOS
MC815	· v	=	=	sec-but	= 5	_	_	<b>=</b>	ວ	183-184	eyclohexane/benzene	55	Civiliacipnios
MC888	S	=	=	C-pent	= 5	_	_	=	ວ	185-186	cyclohexane	54	ChH,CI,N,OS
MC891	S	=	=	c-hex	= 5	_		=	ວ	200-201	cyclohexane/benzene	\$	Chillicinos
MC871	S	=	Ξ	Mc	F =	_	-	=	Ŀ	197-198	benzene	95	CLUINFINOS
MC860	s	=	=	iso-prop	=	_	_	=	Ċ.	174-175	cyclohexane	74	Chillianios
MC872	s	=	=	. pnq-u	±	_		=	Ŀ	126-127	cyclohexane	46	C <sub>IS</sub> II <sub>II</sub> F <sub>I</sub> N <sub>2</sub> OS
MC866	· 05	=	=	iso-but		_	_	=	ŭ.	136-137	cyclohexane	49	CISH,F,N,OS
MC848	· •	=	=	sec-but	=		=	=	Ŀ	149-150	n-hexane/cyclohexane	<del>2</del>	Cr.H.F.N.OS
MC867	· •	: =	=	c-nent	=		=	=	ت	691-891	cyclohexane	45	ChIINFN,OS
MC870	· ·	: =	=	c-hex	. C.	_	=	I	Œ	164-165	cyclohexane	40	CulliFNOS
MC1001	· •	: =	ž	iso-pron	. 5	_	=	=	ວ	196-196.5	cyclohexane/benzene	52	C <sub>t</sub> , II <sub>I</sub> , CI, N, OS
MC996	; 5	: =	Ž	c-nent		_	=	=	ಶ	181-182	cyclohexane	45	C,H,CI,N,OS
MC1016	· •	Ξ.	Ž	c-hex	ַ	_	×	=	ಶ	211-212	cyclohexane/benzene	42	C <sub>I</sub> ,H20CI,N;OS
MC1000	· •	: =	ā	iso-prop	5	_	=	=	ច	166-168	diethyl ether	<b>5</b> 4	C16H <sub>IN</sub> CI <sub>2</sub> N <sub>2</sub> OS
MC1002		: =	iŒ	c-nent	- -	_	=	Ξ	ວ	168-169	diethyl ether	9	C <sub>Is</sub> H <sub>20</sub> Cl <sub>2</sub> N <sub>2</sub> OS
MC1003	· ·	: =	i	c-hex	- 5	_	Ξ	=	ວ	198-199	cyclohexane	<del>-</del> 7	CyllyChyNoS
MC1007		: =	ž	iso-oro	 : <u>:</u>	_	=	=	Ŀ	155-156	cyclohexime	53	C <sub>1</sub> ,11,F,N,OS
MC1044	· •	: =	Me	iso-but	_ 	_	=	=	ᄕ	159-160	cyclohexane	43	Chillin F. N.O.S
MC1045	· 50	=	Σ	n-but	<u>.</u>	_	=	=	Ľ	149-150	cyclohexane	28	C.I.II.F.N.OS
MC1110	· 50	=	Ä	sec-but	<u>-</u>		=	=	œ	133-134	n-hexane	75	C. III.F. N.OS
MC1008	·	=	Ä	c-pent	<u>.</u>	_	=	=	<b>L</b> .	165.5-166.5	cyclohexane	8	C,H,F,N,OS
MC1013	s	=	Ä	c-hex	٤	_	=	¥	Ľ.	206-207	penzenc	77	CINT NEW OS
MC1005	s	=	ជ	iso-prop	<u> </u>	_	=	=	Ŀ	149.150	cyclohexane	<del>\$</del>	ClaH,F,N,OS
MC1006	s:	Ξ	យ័	c-bent	<u>.</u>		=	=	Ŀ	141-143	cyclohexane	<b>.</b>	Challant NiOS
MC1014	တ	=	ជ	c-hex			=	=	<u>a-</u>	154-155	cyclohexane	<u>.</u>	ChillippingOS
MC971	S	=	Ř	iso-prop	CI4=CI1-CH=CI1	=	=	I	I	161-162	n-hexane/cyclohexane	æ :	ClyEnglos
MC972	s	=	ğ	c-pent	C11=C11-C14=C11	<u>~</u>	=	=	=	140-141	n-hexane/cyclohexane		
MC974	S	I	Ψc	c-hex	CII=CH-CII=CI	=	=	=	=	177-178	n-hexane	<b>5</b> ;	Christin
MC969	s	=	ជ	iso-prop	CII=CII-CII=CH	I	×	= :	= :	163-164	cyclohexane	, 4	CULTURAL S
MC973	S	=	ជ	c-pent		=	=	=	Ξ	oi oi		8 ;	C25/14/14/2
MC975	s	=	a	c-hex	CII=CII-CII=CH	I	=	=	=	126-127	n-hexane	<del>-</del> ,	SOLVATION OF
MC844	s:	Me	Ξ	sec-but	Μc	=	=	=	=	177-178	cyclohexane	SS :	Chlippios
MC845	S	ž	×	sec-but	=	=	Ψe	=	=	127-128	n-hexane	<b>.</b>	Chilianics
MC925	S	Mc	=	sec-but		Š,	Ŧ	I	=	163-164	cyclohexane/henzene	æ i	Ciol Nation
MC924	S	Ä	=	sec-but	<b>=</b>	=	ć	=	I	178-180	cyclohexane/benzene	3 9	CipH <sub>14</sub> N <sub>2</sub> O <sub>2</sub> N
MC909	s	Me	=	sec-but	5	=	=	=	=	170-171	cyclohexane	š	Chill recilly OS

l'able 1.		_	Physica	al and Chemica	Physical and Chemical Data of MC Compounds (continued)	mpounds	; (continu	(po				
Compd.	×	>	2	×	R, K;	ĸ	~	ž	m.p., °C	Recryst, Solvent	%. yield	Formula "
• .		:	;	•	5	=	=	=	145-146	cyclohexane	75	ChillyCINiOS
MC'910	S.	NG.	= :	sec-but	==	: 5	: =	: =	163-165	cyclohexane	79	C.II.II.CIN.US
MC911	s.	Š	= :	sec-pat	= =	3 =	: =	: =	170 5-121.5	cyclohexane	65	C. J. I. FN.OS
MC913	ک	ž	= :	sec-but	_ :	= =	= =	: =	146-147	cyclohexane	ננ	C.L.II.FIN,OS
NC918	S.	ğ	=	sec-but	= :	<b>-</b> :	= =	= =	154.155	cyclohexine	69	SO,N:1,,11,,13
MC'919	×	ž	=	sec-hu	= 1	= :	= =	= 5	196-306	binzig:	5	C., H., CI, N, OS
MC:912	s	Ϋ́с	=	Me	= : :	= :	= :	3 5	107-007	mulchus man Assesser	, ×	CLICINOS
MC914	s:	Σ	=	iso-prop	= <sup>-</sup>	= :	= :	5 5	747-147	Cyclinicaline Control Control	Ç	CHUCHNOS
MC920	s:	Me	=	n-hut	= 5	= :	= :	3 5	001-6/1	Cyclothagus	18	C. II. CI,NOS
MC'916	s	Ψe	=	iso-bul	= 5	= :	= :	J :	607-907	Cyclimeranic	5	C. II. CI,NOS
MC850	s	ž	=	sec-but	= : = :	= :	= :	J (	507-507	cyclohexanellenzene	€	C. H. CI.N.OS
MC915	s	χ̈	=	c-pent	= : ::	= :	= :	3 5	056-767	cyclinicality concerns	. <del>4</del>	CHCANOS
MC917	s	Ř	=	c-hex	= : 	= :	= :	ء ت	007-107	Lycharica mic	ć,	SO,N.F. II.
MC869	s	Ä	=	Mc	<u>-</u> :	= :	= :	L	210.3-217.3	control	92	C. H. I'N OS
MC881	s	ğ	=	iso-prop	<b>=</b> :	= :	= :	<u>.</u> [	071 071	cyclohexane	59	C., H., F., N, 0S
MC905	ss	Ä	=	n-but	=	= :	= :	<u>.</u> (	671-071	Cyclothorune	2	C. II. F.N.OS
MC921	s	Ä	Ξ	iso-but	=	=	=	<u>-</u> 1	791-191	cyclonexime	Ç Ç	SO'N'E E
MC849	· 22	Me	Ξ	scc-hut	<b>=</b>	=	=	Œ.	128-129	n-nexane	, .	SCINE HILL
MCSOT	: 0	ž	=	c-nent	=======================================	=	=	ᇆ	192-193	cyclohexanc	94	
MC.922	n 0	, , ,	: =		. E	=	=	Œ.	191-192	cyclohexane	67	CO.N. J. C. L.
MC.92.3	n :	2 2	: 2	M.	. 2	=	Ξ	뜨	202-203	eyelohexane/henzene	<del>ن</del> تا	CONTRACT
MCIOSO	n :	2 2	2 2	rac had	. 12	=	=	Œ	135-136	cycluhexane	55	Culting
MC.1109	n s	2 2	2 2	Sec-Dui	. 2	=	=	ت	196-197	cyclohexane	9	CIRTINOS
MC104/	<u>ر</u> :	<u>د</u> ک	ž :	c-pent	. =	=	=	<b></b>	140-141	n-hexane	47	ChluNoS
MC798	<u>بر</u>	<b>i</b>	= :		: =	: <b>=</b>	=	Œ	174-175	benzene	82	Chally F.N.OS
MC1037	s :	<u>ವ</u> (	= :	Iso-prop	. 2	: =	: =	. <u>:</u> -	150-151	n-hexane/eyclohexane	G	CIMENTOS
MC1038	'n	១ :	= :	sec-pni	11.7-11.7 11.7-11.7	: =	: =	=	198,5-199,5	cyclohexane	45	Cillianos
MC804	×	酉	= :	scc-but		= =	: =	: :	167-168	n-hexane	76	CI-II FINOS
MC1039	s	i-pro	=	iso-prop		= =	= =	. =	127 5-128 5	evelohexane	89	C.H.N.OS
MC852	s.	allyl	=	sec-but	= :	= =	= =	= =	108-100	n-hexane	42	C,N,H,N,OS
MC856	v:	n-pro	=	sec-hut	= :	= =	= =	: =	: : : :		32	C.L.II.N.OS
MC834	S	n-bat	=	sec-but	= :	= =	= =	: 2	138-140	n-hexane/evelohexane	20	ChilliPiNO
MC:1119	Ī	=	= :	cthyl		= =	=	. Œ	136-137	cyclohexane	49	CHIRENO
MC1078	Ξ	= :	= :	n-prop	ء ۔	: = 	: =	٠ نـ	150-151	diethyl ether	28	C, JI, F, N,O
MC979	Ī	=	= :	iso-prop		= =	: =	. <i>u</i>	183-184	eyclohexane/Denzene	89	ChilliFiNO
MC980	Ī	=	= :	c-brop		= =	=	. (2	130-131	n-hexane	9	C, H, F, N,O
MC1077	Ī	Ξ	I	n-hut			: =	. (z	140-141	diethyl ether	8	CLUFINO
MC945	Ē	Ξ	=	sec-but	- :	= =	= =	. 0	120-121	acetonitrile	78	C,H,F,N,O,
MC1043	Ē	Ξ	Ξ	McOethyl	 - :	c =	= =	. 11	: ::=		74	Cl,H1,F2N,O
MC1022	Ξ	=	I	c-bcut	- ·	= =	: =	. µ	143-144	diethyl ether	45	ChHIFING
MC1049	Ī	=	=		<u>.</u>	= =	= =	. (1	: : :		48	C,111,F2N,O
MC1048	Ē	=	ž	•	· :	= =	= =	- 6	165.166	n-hexane	53	CHIPFINO
MC1118	Ē	Σ	=	iso-prop	۔ د	= :	= =	<u>.</u> [	107-100		Š	CloH, F, N,O
MC1130	Ī	ğ	=	sec-but	ii. i	= =	= =	2 (2	115-117	n-hexane/eyclohexane	9	C,H,F,N,O
MC1050	Ī	Σ	<b>=</b> :	luad-o		= =	: =	. 12	182-183	cyclohexane/benzene	82	C <sub>IN</sub> II <sub>I</sub> ,F,N,O
MC1105	Ξ	ž	Ĩ	benzyl	<u>-</u>	= =	:	•				

Table 1.			Physica	Physical and Chemical Data of MC Compounds (continued)	I Data of MC	Comp	onnds (c	ontinuc	<del>-</del>				
Compd.	×	>	2	~	۲.	괃	ş≃	7≃	ž.	ո.թ., ° С	Recryst. Solvent	%. yield	Formula "
		2	ž		£	=	=	=	:-	io	:	38	C,N,F,N,O
MC.1129	Ē :	ž :	ž :		. 2	: =	: =	: =	<u>د</u>	202-203	acetonitrile	36	CLNFIND
MC1167	Z	Ξ	= .	Z c	<u>.</u> (	= :	: :	: :	. 6	210.211	postonitrile	48	C'II'E'N'O
MC1168	Ē	ž	=	Ä	<u>.</u>	=	= :	= :	L (	117-017		: 3	CNIII
MC1186	Ī	ž	Ξ	ս-եւսր	Œ,	=	=	=	<u>*</u>	/61-96	nceromitine i	1 3	
MCHRS	Ī	Ä	Ξ	n-pnt	Œ.	=	Ξ	=	Ľ	192-193	acetonifrie	e ·	Challering
MC1178	ž	=	Me	Me	Ľ	=	=	x	ᄕ	145-146	acetonitrile	75	ChillippyO
ON COLUMN	Ž	: =	ž	ם-סנסים	· CL	=	=	=	뜨	oi	•	45	CisH,F,N,O
MC1130	Ž	: =	į	iso-oron	٠. د	=	=	=	ت	oil	;	54	Civili,Fin,O
MCIIS	Ž	: =	Ž	a-hut	. "	=	=	=	Œ	oit	:	55	CleHyFyN,O
MC1163	. ž	= =	Ž	s.chut	. "	=	=	=	Ľ	oil	:	50	C, 11, F, N, O
MC1192	2	: =	Ž	c-hev	. 12	: =	=	=	Œ	io	1	62	C,NF,N,C
MC1180	2 2	: X		Me	. L	=	=	=	Ľ.	193-194	eyelohexane/henzene	74	C, H, F, N, O
2012	2	2 2		, in the	. "	=	=	=	Œ	oi:		49	Cinfilling
		2 2		100-100	. Ľ	=	=	=	Ĺ	ej.	:	54	C <sub>l</sub> ,H <sub>2</sub> ,F <sub>2</sub> N <sub>3</sub> O
MCIIO	Ē 2	ž		Z	. tr	=	=	Ξ	Ľ.	210-211	cyclohexane/henzene	88	C,JH,F,NO
MC182	z 2	= =		Me singer	. "	: =	: =	=	Ľ	195-196	acetonitrite	84	ChH,FJN,O
MCIISS	2 2	= =	-	morph	. tı	: =	=	=	Œ	215-216	acetonitrile	75	CIAHISFINIO;
MCII88	z 2	= =		thiomorph	. <u>(</u>	: =	: =	=	뜨	233-234	acetonitrile	78	C.H.F.N.OS
WC175	z 2	= =		miconico pui piporid	. c	: =	: =	<b>=</b>	Ľ.	209-210	acctonitrile	89	C,N,F,N,C
* 10 E	2 2	= =		pilonid	. 12	Ξ	=	=	Œ	233-234	acctonitrile	22	CISHISF3N3O
MC1363	z 2	= =	: =	Francisco (Control of Control of	. د	=	=	=	ï	159-160	acetonitrile	43	C <sub>IS</sub> III7F <sub>3</sub> N <sub>3</sub> O
MC1202	2 2	= =			. (:	: =	=	Ξ	Œ	111-112	n-hexane	35	CITHILENO
MC1204	z :	= 3	= =	rahand-m	نا ـ	: =	: =	=	Œ	237-238	acetonitrile	80	C, III, F, N, O
MC1195	z	Σ		Mc <sub>2</sub>	. L	: =	: =	: =	. 4	235-236	acetonitrile	62	C <sub>1</sub> ,H <sub>2</sub> ,F <sub>2</sub> N <sub>4</sub> O
MC1203	z 2	Z Z	= =	morph	. tr	: =	: =	=	ᄕ	244-245	acetonitrile	65	C <sub>th</sub> H <sub>1</sub> ,F <sub>2</sub> N <sub>3</sub> O <sub>2</sub>
MCIZIO	2 2	2 3	: =	thiomorph	. "	: =	=	Ξ	Œ	255-256	acetonitrile	24	C.L.I.,F.N.OS
MC1200	2 (	<u> </u>		in peop	. Œ	: =	=	=	٠	177-178	n-hexane/cyclohexane	45	C.H.F.N.OS
MC11.57	<u>ر</u>	ž :		180-prop		: =	: =	: =	. 4	122-123	n-bexane	15	C., IL, F, N, OS
MCII /S	<u>ر</u>	Ξ:		100-11	نا خ	: =	: =	=	(*	152-153	cyclohexane	28	C, II, F, N, OS
MC115.3	n :	ž :	ΣZ	180-0111	L (I	: =	: =	: =	<u>ن</u>	208-209	n-hexanc/eyclohexane	84	C <sub>1</sub> ,H <sub>22</sub> F <sub>2</sub> N <sub>2</sub> OS
MC11/4	n:	ž:		C-nex		: =	: =	: =	<u> (1</u>	159-160	cyclohexane/benzene	72	C <sub>IJ</sub> H <sub>IJ</sub> F <sub>2</sub> N <sub>2</sub> OS <sub>2</sub>
MCII61	n e	= :		Mesime	. "	: =	: =	: =	4.	183-184	cyclohexane/benzene	20	CITHE NOS
MC1162	'n	¥ W		Mesme	ا نا	: =	: =	: =	, <u>t</u>	153-154	cyclohexane	69	C <sub>13</sub> H <sub>16</sub> F <sub>2</sub> N <sub>2</sub> OS <sub>2</sub>
MC1157	s o	<b>ញ</b> .		McSMe	L	= =	: =	: <b>=</b>	. LT.	158.5-160	cyclohexane	62	CleHINF,N,OS,
MC1145	S)	-pro	<b>z</b> :	Mesme	L 2	: 3	: =	: =	. =	117.5-118	n-hexane	64	Culli,N2OS2
MC1140	တ	I		Mesme	5	5	:	:	:				

\*All compounds were analyzed for C, H, N, S, and, when required, Cl and F; analytical results were within ±0.4% of theroretical values.

	, IS	40	6	>6.7	36	25	<u> </u>	<b>V</b>	, ,	777<	232	248	250	>200	>154	= <del>-</del>	\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	333.3	>800	392	101	7 2 3	ž č	÷ ;	60/	>280	5,	V	<b>%</b> ,
		3.5	6.4	30	3.5	25	50	ξ,	×01	<i>zi ,</i>	e ,	ė:	∞i ¦	1.0	1.3	æ: ;	3.4	9.0	0.25	0.40	<u>s:</u> .	- (	7 1	Λě	0.26	0.7	8.7	21.2	23
	СС <sub>50</sub> , [µМ]	143	28	>200	138	130	>200	>200	19	>200	159	149	200	>200	>200	>200	>200	200	>200	157	151	200	911	071	200	>200	>200	>200	>200
	ž	Ξ	Ξ	=	=	<u> </u>	<u>.</u>	= :	= :	= :	<b>=</b> :	=	Ξ	=	=	=	=	=	=	=	<b>=</b> :	= :	= :	<b>=</b> :	Ξ	Ξ	=	=	=
	~≃	Ξ	=	Ξ	Ξ	=	=	= :	=	= :	=	=	Ξ	=	=	=	=	=	=	=	=	=		<b>=</b> :		=	Ξ	=	<b>=</b>
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3	R²	=	=	=	Ξ	=	<b>=</b>	Ξ	Ξ	=	I	Ξ	Ξ	H	Ξ	=	<u></u>	=	H	Š N	=	Ξ	ರ	I	=	Ľ	I	=	Ξ
æ e	<u>~</u>	, Ħ	Ξ	Ξ	Ξ	Ľ	ᄄ	=	=	Ξ	I	=	H	I	H	Mc	Mc	Ξ	NO,	Ξ	H	ರ	=	Ξ	Ľ	¥	=	ZHZ	Ξ
MC Compounds.	· <b>జ</b>	2,5-Mc,-c-hcx	4,5-Me <sub>2</sub> -c-hex	3,5-Me2-c-hex	2,5-Mc2-c-licx	sec-but	c-pent	henzoyloxymethyl	sec-but	iso-prop	c-pent	c-hex	iso-prop	c-pent	c-hex	sec-but	c-pent	sec-but	sec-hut	sec-hut	scc-but	sec-but	sec-but	sec-but	sec-but	sec-put	scc-hut	sec-but	sec-but
	7	×	=	Ξ	Ξ	Ξ	Ξ	Ξ	Ph	Mc	Me	Mc	酉	ភ	ជ	H	Ξ	I	=	Ξ	I	Ξ	Ξ	Ξ	=	Ξ	=	Ξ	Ξ
unti-HIV-1 Act	>-	I	I	=	Mc	H	=	Ξ	Ξ	<b>=</b>	H	Ξ	=	=	Ξ	-	=	===	: =	Œ	H	H	H	=	Ξ	=	H	=	H
oxicity and	×	0	0	0	0	0	0	S	S	S	S	S	S	S	v.	S	· ·	S	v.	S	S	S	S	S	S	S	S	S	S
Table 2. Cytotoxicity and anti-HIV-1 Activity of	Compd.	MC 507	MC 508	MC 512	MC 531	MC 1114	MC 1103	MC 843	MC 796	MC 890	MC 892	MC 898	MC 899	MC 900	MC 903	MC 806	MC 842	MC 809	MC 817	MC 897	MC 863	MC 854	MC 857	MC 859	MC 880	MC 884	MC 889	MC 825	MC 960

Table 2. Cyt	Cytotoxicity and anti-IIIV-1 Activity of	ınti-IIIV-1 /	Activity of	MC Compounds (continued)	nucd)			<b>.</b>				
Compet	>	>	7	~	, W	κ,			χ.	[htM]		»IS
comba.	•	•	1	:						CC.,,	- 201	,
MC 868	S	=	=	sec-but	CF,	=			= :	>200	32	0.2 0
MC 959	S	Ξ	=	sec-but	=	=			=	200	25	œ ,
MC 952	S	=	=	sec-but	OMc	=				>200	1.96	>208
MC 957	S	=	=	sec-but	=	OMc				>200	1:2	99 <b>I</b> <
MC 964	တ	=	=	sec-but	=	=				147	4	10.5
MC 1041	S	=	=	sec-but	=	<u>-</u> -				>200	4.	>143
MC 1042	v	=	=	sec-but	Ξ	Me				133	9.0	222
MC 877	S	=	=	Mc	כ	=				>200	3.2	>62
MC 878	S	=	=	iso-prop	ฮ	=				>200	1.9	>105
MC 886	S	=	Ξ	n-but	ס	Ξ				>200	0.44	>454
MC 885	· v:	Ξ	Ξ	iso-but	ಶ	I				>200	0.45	>444
MC 815	y v	: =	=	sec-but	ರ	=				>200	0.14	>1,428
888 CW	o vo	: =	=	c-pent	ū	=				>200	0.4	>200
MC 891	v.	: =	=	c-hex	ט	=				>200	9.0	>333
MC 871	o vo	: =	=	Mc	ت	Ξ				200	0.81	247
MC 860	o on	: =	=	iso-prop	ᄄ	I				>200	0.2	×1,000
MC 877	y V	: =	: =	n-but	Ľ	=				162	0.18	006
7/8 CW	, v	: =	: =	iso-but	ٺ	=				182	0.14	1,300
MC 848	) V.	: =	=	sec-but	Ľ					200	0.04	5,000
MC 867	o v	: =	: =	c-pent	1	=				>200	0.08	>2,500
MC 870	o v	: =	: =	c-hex	Ľ	I			<u></u>	200	0.08	2,500
MC 1001	o 01	: =	Me	iso-prop	ប	=				117	1.2	97.5
MC 996	o vo	=	ğ	c-pent	ס	=				78.3	0.1	78.3
MC 1016	v	Ξ	Mc	c-hex	ರ	Ξ				>200	2.9	69×
MC 1000	o vo	: =	ច	iso-prop	כ	Ξ				>200	0.4	>500
MC 1002	S	=	<u></u>	c-pent	ರ	=				23.4	0.1	23.4
MC 1003	S	<u></u>	酉	c-hex	ರ	<b>=</b>				>200	3.6	5.5.5
MC 1007	S	=	Me	iso-prop	Ċ.	=				191	0.05	3,340
MC 1044	· v	-	Me	iso-but	ï	=				>200	0.05	>4,000
	o Co	: =	Mc	n-but	متا	=				>200	0.02	2,857
MC 1110	o co	=	Me	sec-hut	ᄕ	H				>200	0.03	999'9<
	S	=	Mc	c-pent	÷	=				>200	0.03	>6,666
	S	Ξ	Me	c-hex	ប	Ħ		=	Œ.	>200	0.16	052,1<
	S	Ξ	酉	iso-prop	ᄄ	=	=	=	ii.	92	90.0	6/8
MC 1006	S	=	ជ	c-pent	<u>ب</u>	=	=	=	<u></u>	200	0.15	1,555

Ŝ	Cytotoxicity and anti-fit v-1 Activity of X Y Z	×		<u>"</u> ഷ	-⊱	ž	[Fig		, IS
១	o-pe	*	: :		=	ŭ.	130 0.		2,600
Mc	iso-pr	dο	CII=CII-CII=CII		=	= :		<del>-</del> . '	80 !
	c-pent			= =	= =	= =		<u>د</u> <u>۲</u>	180
Me G	c-hex	,			= =	= =		<u>+</u> ~	33.3
ថេ	c-pent	_	CII=CII-CII=CH		: =	: =		9 0.	17
Ö	c-hcx		CH=CH-CH=CH		=	=		<u>∞</u>	94
Ħ	sec-hut		Mc		Ξ	=		1.7	×118
=	sec-but		H		<b>=</b>	=		æ: ˈ	32
Ξ	sec-put		11 NO;		=	=		.35	>571
H	scc-pat		11	_	=	Ξ :		C1	×100 ×100
=	sec-but		= 5		=	=		.27	>74
=	sec-but				=	=		96.	>208
Ξ	sec-but		]]		=	=		5.5	23
=	sec-but		= 4		=	=		14.	341
=	sec-but		<u> </u>		=	=		<u> </u>	>166
=	sec-but		= ==		=	=		=	9.5
Ξ	Me		<b>=</b> 5		=	ರ		3.2	<b>&gt;</b> 05
Ξ	iso-prop		<b>□</b>		Ξ	ರ		: : ۲	>154 :
Ξ	n-but	>	C H		=	ت ت		.17	Z :
13	iso-but		5		Ξ:	บ :		<u>.</u> 2	>166
=	sec-put		T U		=	ට :		.05	×,000
=	c-pent		C C		=	ರ		œ. ;	<u> </u>
Ξ	c-hcx		<u> </u>		Ξ	ប		22	ر م
=	Mc		<b>=</b>		=	<b>:</b>		61.	1,053
11	iso-prop		<b>1</b>		=	<u>:</u>		.05	×4,000
=	n-but				=	ű.		.0 <del>.</del>	>2,500
Ξ	iso-but		F H		=	<u>-</u>		0.1	640
H	sec-but		F		=	Ľ,		100.	8,000
Ξ	c-pent		F -		=	ت		80.0	>2,500
	c-hcx		H 4		Ξ	<u>-</u>		.09	>2,222
Me	Me		F .	=	=	뜨		.04	>5,000
Me	sec-but		F	¥	=	Ľ.		.03	999'9
Mc	c-pent		F	=	=	<u>~</u>		600.	>22,222
=	sec-but		=======================================	=	=	Ξ		0.1	>200

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	æ	=	=	=	=	Ħ	I	=	Ξ	=	=	=	Ξ	=	=	=	=	=	=	=	=	=	=	=	=	=	=	=	=	=	=	=	Ξ	Ξ	=
	5≃	=	=	-CII=CII	Ξ	=	=	H	=	=	=	I	Ξ	=	I	=	=	<b>=</b>		=	I	I	=	=	Η	=	=	=	=	=	=	=	=	H	Ξ
ontinucd)	R.	<u>(•</u>	. <u>:</u> -	CII=CII-CI	<u>(:,</u>	=	=	11	Ľ.	Ľ.	Œ,	Ľ	뜨	<u></u>	Ľ	ت	ᄄ	ഥ	ű.	ഥ	<u></u>	ᄕᅩ	Ľ	ت	Ľ	Ľ.	Ľ,	<u>-</u>	뜨	Œ	<u>-</u>	(Ľ	뜨	Œ,	Ľ
of MC Compounds (continued)	~	or sign	sec-but	sec-but	iso-prop	sec-but	scc-but	sec-but	cthyl	n-prop	iso-prop	c-prop	n-but	sec-put	McOcthyl	c-pent	c-hex	c-pent	iso-prop	sec-but	c-pent	benzyl	c-pent	Mc	Mc	n-prop	n-but	Mc	n-prop	iso-prop	n-but	sec-but	c-hex	Mc	p-but
	2	2		=	=	=	=	Ξ	<b>=</b>	Ξ	I	H	H	=	=	Ξ	I	Me	Η	I	I	Ξ	H	=	Œ	=	Ξ	Ψ¢	Me	Mc	Me	Mc	Mc	Mc	Ž
Cytotoxicity and anti-IIIV-1 Activity	>	2	ថេ	១	iso-prop	allyl	n-prop	n-but	H	H	H	H	H	=	Ξ	=	I	==	Me	Me	Mc	Mc	Mc	H	Mc	Me	Mc	Ξ	=	=	I	H	Ξ	Mc	Me
totoxicity an	×	υ	o v	တ	S	S	S	S	Ē	Ξ	Ξ	ΞŽ	Ī	ž	Ē	Ξ	Ē	Ī	Z	Ē	Ī	Ī	Ē	Ē	Ξ	ΞZ	Z	Z	ΞZ	Ē	Ë	Ē	Z	H	Z
Table 2. Cyt	Compd.	MC. 1037	MC 1038	MC 804	MC 1039	MC 852	MC 856	MC 834	MC 1119	MC 1078	MC 979	MC 980	MC 1077	MC 945	MC 1043	MC 1022	MC 1049	MC 1048	MC 1118	MC 1130	MC 1050	MC 1105	MC 1129	MC 1167	MC 1168	MC 1186	MC 1185	MC 1178	MC 1190	MC 1191	MC 1189	MC 1192	MC 1180	MC 1170	MC 1187

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'	اري. م مو	CO:0	7.1	9.0	0.05	0.02	2.1	0.26	3.8	0.02	0.36	0.047	0.09	0.007	0.008	0.01	0.018	0.002	0.7	0.80	0.12	0.11	0.10	20	
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Compd.		MC 1182	MC 1183	MC 1188	MC 1193	MC 1194	MC 1196	MC 1202	MC 1204	MC 1195	MC 1203	MC 1205	MC 1206	MC 1137	MC 1175	MC 1153	MC 1174	MC 1047+	MC 1047-	MC 1161	MC 1162	MC 1157	MC 1145	MC 1140	

" Compound dose required to reduce the viability of mock-infected cells by 50%, as determined by the MMT method. Compound dose required to achieve 5t1% protection of MT-4 cells from HIV-1 induced cytopathogenicity, as determined su/EC so ratio. " Data represent mean values of at least two separate experiments. " Selectivity index, CC by the MTT method.